Conscious and unconscious vision

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Chapter 2.

Masking disrupts recurrent processing in human visual cortex

Abstract

In masking, a stimulus is rendered invisible through the presentation of a second stimulus shortly after the first. Over the years, authors have typically explained masking by postulating some early disruption process. In these feed forward type explanations, the mask somehow “catches up” with the target stimulus, disrupting its processing either through lateral or interchannel inhibition. However, studies from recent years indicate that visual perception - and most notably visual awareness itself – may depend strongly on cortico-cortical feedback connections from higher to lower visual areas. This has led some researchers to propose that masking derives its effectiveness from selectively interrupting these recurrent processes. In this experiment, we used EEG measurements to determine what happens in human visual cortex during detection of a texture defined square under non-masked (seen) and masked (unseen) conditions. EEG derivatives that are typically associated with recurrent processing turn out to be absent in the masked condition. Moreover, extrastriate visual areas are still activated early on by both seen and unseen stimuli, as shown by scalp surface Laplacian current source density maps. This conclusively shows that feedforward processing is preserved, even when subject performance is at chance as determined by objective measures. From these results we conclude that masking derives its effectiveness, at least partly, from disrupting recurrent processing, thereby interfering with the neural mechanisms of figure-ground segregation and visual awareness itself.

Introduction

In backward masking, a target stimulus is rendered less perceptible or even invisible through the presentation of a second stimulus, the mask. By now, a respectable number of studies have investigated the neural basis of visual masking, both in its own right, and as a corollary to studying visual perception and awareness. In explaining a variety of masking phenomena, some have emphasized low level lateral inhibitory mechanisms (Francis, 1997; Macknik & Livingstone, 1998), while others have postulated target disruption at the pre-categorical level through interchannel inhibition (Breitmeyer, Ro, & Ogmen, 2004; Ogmen, Breitmeyer, & Melvin, 2003).

However, early studies as well as more recent imaging studies on masked priming show that processing of masked targets may continue well beyond the pre-categorical level (Dehaene, et al., 2001; Dehaene, et al., 1998; Eimer & Schlaghecken, 1998; Greenwald, Klinger, & Liu, 1989; Naccache, et al., 2005). These results suggest that masking does not disrupt target processing at an early stage. This is in line with various neurophysiological reports in which early transient responses to undetected masked stimuli were measured in high visual areas such as IT, and even in visual neurons of the frontal eye field (G. Kovacs, et al., 1995; Rolls & Tovee, 1994; Thompson & Schall, 1999).

This has resulted in a number of researchers proposing that masking disrupts reentrant cortical signals, while largely leaving signals related to feedforward processing intact (Bridgeman, 1980; Di Lollo, et al., 2000; Lamme, et al., 2002; Ro, Breitmeyer, Burton, Singhal, & Lane, 2003). These proposals build on the notion that besides feedforward connections from low to higher visual areas, roughly equal numbers of recurrent or reentrant connections exist going back from higher to lower visual areas (Felleman & Van Essen, 1991). Such feedback connections have been suggested to play an integral role in a range of processes, such as figure-ground segregation, attention, awareness, predictive coding, as well as perception of visual detail (Hochstein & Ahissar, 2002; Kastner & Ungerleider, 2000; Lamme, 1995; Lamme & Roelfsena, 2000; Luck, Chelazzi, Hillyard, & Desimone, 1997; Rao & Ballard, 1999; Spratling & Johnson, 2004). It is not unlikely that masking derives its effectiveness (at least in part)
from disrupting recurrent signals, while leaving feedforward signals intact, thereby interfering with mechanisms thought to mediate visual awareness itself (Lamme & Roelfsema, 2000).

However, many masking studies do not show a relation between recurrent processing and masking. This may be caused by the fact that many such studies make use of luminance or color contrast targets and metacontrast masks. In such metacontrast masking paradigms, the mask tightly fits the preceding target and the common contour has a contrast of opposite polarity (Becker & Anstis, 2004). It is quite likely that low level lateral inhibitory mechanisms come into play when processing these types of stimuli, at least at sufficiently short SOA’s (Macknik & Livingstone, 1998; Macknik & Martinez-Conde, 2004).

Contrast sensitivity is fundamental to vision and contrast is greatly enhanced by lateral inhibition at many levels of visual processing, starting in the retina (Hartline, 1949). Therefore, the use of (meta)contrast stimuli without appropriately controlling for the influence of contrast on the signal exaggerates the influence of low level lateral inhibition, while obscuring the influence of other higher level processes, such as recurrent processing. The stimuli used in this experiment - in combination with a subtraction method (detailed below) – eliminate the influence of luminance contrast on the signal, while at the same time controlling for other low level influences.

Aside from the contrast issue, visibility of masked targets is often assessed using an identification task. However, a detection task is more fundamentally connected to the notion of visual awareness, as many studies demonstrate that residual identification in the absence of awareness exists. This is evident not only from research into the phenomenon of blindsight (Weiskrantz, Barbur, & Sahraie, 1995; Weiskrantz, Warrington, Sanders, & Marshall, 1974), but selective behavior in the absence of awareness has also been shown in normal subjects (Boyer, Harrison, & Ro, 2005; VanRullen & Koch, 2003) as well as in studies using masked priming (e.g. Eimer & Schlaghecken, 1998; Vorberg, Mattler, Heinecke, Schmidt, & Schwarzbach, 2003). Selectivity thus seems an inappropriate indicator of awareness. Therefore, the present experiment employs an objective detection task to assess visibility.
We investigated the effect of pattern masking on the detection of an orientation defined square in a texture (Figure 2.1), while recording EEG in human subjects. Figure and No Figure trials were either strongly masked by a pattern mask leading to chance detection performance (Masked condition), or weakly masked by a uniform grey screen leading to near perfect detection performance (Unmasked condition). We used a subtraction procedure to control for various confounding influences on the EEG.

**Figure 2.1 Schematic description of a trial.** Subjects were required to indicate whether a figure was present in a texture or not. Half of the trials contained a figure. Half of the trials only contained a homogenous texture. A trial could either be strongly masked (masked) or weakly masked (unmasked). If subjects could not see anything due to masking they were instructed to guess (forced choice detection).

By subtracting the Figure trials from the No Figure trials (see Figure 2.2), two goals were achieved: (1) because both Figure and No Figure trials were followed by a mask in the masked condition, any direct influence of the mask on the EEG was subtracted out. The same holds in the unmasked condition. (2) Because both Figure and No Figure trials were created using the same set of oriented line elements, Figure and No Figure trials contained exactly equal amounts of contrast and orientation. Thus, any direct influence of low level processes such as contrast detection and orientation tuning is subtracted out (Zipser, et al., 1996). This subtraction procedure would not work for metacontrast target-mask combinations as used in other studies, as the net result of the subtraction of target-present and target-absent trials would inherently leave a net result of contrast.
Figure 2.2 (a) Examples of stimuli and the schematic orientations scheme. Each grey value represents an orientation, white is isoluminant grey. (b) Representation of subtractions leading to correlates of global processing. Influences of local line elements as well as direct mask contributions are canceled out by ensuring each orientation is present equally often at each side of the minus sign. Color scheme same as in (a).
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The spatiotemporal dynamics of the target-present and target-absent subtraction signal was studied at an extremely short SOA (16.6 ms) where targets were either easily perceived or remained totally undetected as determined by objective measures. We observed marked activation in extrastriate regions in both the visible and invisible condition as a result of feedforward processing. Later and more posterior activation resulting from recurrent processing was observed only in the visible condition. From the results we infer that recurrent processing is abolished as detection performance drops to chance, even though feedforward processing is entirely preserved.

Methods

Participants
Twenty one Psychology students took part in the experiment in partial fulfillment of first year course requirements. All subjects (mean age 21.4, ±1.9) had normal or corrected-to-normal vision. 71% was right handed. Each subject provided written informed consent before the start of the experiment. All procedures were approved by the ethical committee of the psychology department of the University of Amsterdam. Two subjects scored more than two standard deviations below the mean in the unmasked task, indicating difficulty performing the task properly. These were excluded from further analyses. Three subjects scored significantly above chance in the masked condition as ascertained by binomial tests (p<.05), indicating an ability (however slight) to detect the masked figure at this masking interval. These were excluded because it cannot be ruled out that they were still consciously processing these stimuli. In total five subjects were excluded.

Stimuli and Task
Without prior training, subjects had to detect the presence or absence of an orientation defined square in a texture pattern, which was either followed by a mask or by an isoluminant grey screen (all stimuli had a
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luminance of 72.7 cd/m²). Each of the 4 resulting trial types was presented an equal number of 300 times in random order throughout the experiment. A trial started with 300 ms fixation, followed by a centrally located target stimulus for 16.6 ms, immediately followed by either a mask or an isoluminant grey screen for 500 ms (Figure 2.1). Target stimuli and mask consisted of a texture of oriented line elements (approx. 0.07° thick and 0.4° long in visual angle). All line elements in a target either had the same orientation (homogenous trials), or contained a centrally located square with line elements orthogonal to the background (Figure trials). The square subtended 2.73° of visual angle. In the masked condition, targets were immediately followed by a texture defined pattern mask (4.54° visual angle) using orientations not used in the preceding target. In the unmasked condition, targets were followed by an isoluminant grey screen (no texture). A total of four orientations of line segments were used for creating figure, homogenous and mask stimuli (22.5°, 67.5°, 112.5° and 157.5°, also see Figure 2.2), so that different orientations for target and mask could always be used within any one trial. All orientations were completely counterbalanced over trials so figure and homogenous trials did not differ from each other with respect to local stimulation (for a similar procedure see Caputo & Casco, 1999; Lamme, Van Dijk, & Spekreijse, 1992; Scholte, et al., 2006). Stimuli were created using Matlab (The MathWorks, Inc., Natick, MA, USA). Stimuli were presented using Presentation (Neurobehavioral Systems, Inc., Albany, CA, USA).

Subjects were instructed to fixate throughout the experiment. The fixation dot would turn green 500 ms after target offset, to indicate a response was required, after which the subject had a maximum of 1000 ms to respond before the experiment would continue. On half of the trials subjects were required to press the left button to indicate square detection and the right button for a homogenous trial. In the other half of the trials subjects had to switch so that a square was associated with right and homogenous with left button presses. This response scheme was counterbalanced across subjects, half of the subjects starting out square detection with their right hand, and half of the subjects starting out using their left hand. Switching hands posed no difficulty as the high percentages correct on unmasked trials shows. Subjects were encouraged to guess if they
were unable to see if a figure was presented or not, effectively turning the task into a forced choice task.

**Measurements and Analyses**

ERP’s were recorded from the scalp using a BioSemi ActiveTwo 48 channel active EEG system (BioSemi, Amsterdam, the Netherlands) at 256 Hz. 48 scalp electrodes were measured (referenced to Fz), as well as two electrodes for horizontal and two for vertical eye movements (each referenced to its opposite counterpart). The data was filtered using a high pass filter of 1 Hz and a low pass filter of 20 Hz, as well as a 50 Hz notch filter. Before ocular correction, automatic artefact rejection was applied by removing segments containing voltage steps of more than 50 μV, removing any segments falling outside the -200 μV to 200 μV range as well as removing those segments containing larger than 300 μV differences within the segment. Ocular correction was applied on the basis of the horizontal and vertical electro-oculograms (Gratton, Coles, & Donchin, 1983). After ocular correction, artefact rejection was applied again by removing all segments outside the -75 μV to 75 μV range. Linear local DC detrending was applied. This is a procedure to remove current drift by subtracting a linear function from each segment. Baseline correction was applied in the -300 ms to 0 ms interval. All pre-processing steps were done using Brain Vision Analyzer (Brain Products GmbH, Munich, Germany).

All analyses were done on difference waves. These were obtained by subtracting average No Figure trials from Figure trials, separately for the Masked and Unmasked condition. Spline Laplacian distribution maps were calculated by interpolating difference waves using spherical splines and approximating current source densities (Perrin, Pernier, Bertrand, & Echallier, 1989). The resulting maps are spatial second order derivatives of the scalp potentials lending greater weight to local contributions of cortical generators, filtering out deep sources, as well as being reference free (Nunez & Srinivasan, 2006).

In order to test at which time points the differences between Figure and No Figure trials for the Masked and Unmasked condition significantly deflect from chance, as well as test the difference between masked and unmasked trials, a random effects analysis was...
performed by employing a paired two-tailed t-test at each time point, treating the average of each subject at that time point as an observation. Correction for multiple comparisons with respect to the number of time points being tested was done by limiting the False Discovery Rate (FDR), a method in which the p-value at which significance is evaluated is corrected for the number of tests being performed (Benjamini & Hochberg, 1995). The method fixes the expected proportion of false positives (the FDR) in contrast to Bonferroni correction, which controls the chance of any false positive among all tests. The FDR is an often used method in a wide variety of scientific fields. For an explanation of how it is used in the field of neuroimaging see Genovese, Lazar & Nichols (2002). Statistical analyses and visualization of the timecourses were done using Matlab (The MathWorks, Inc., Natick, MA, USA).

**Results**

The difference between Figure and No Figure trials was easily detected when no mask was present, as expressed in an average detection rate of 96.5% (±2.5). For the masked trials the detection rate was at chance at 51.4% (±4.1). Detection rates are expressed as the average perfect observer score, which is an objective and subject bias free measure of a subjects’ ability to detect a signal (Wickens, 2002). None of the included subjects scored significantly above chance in the masked condition (binomial tests, significance evaluated at .05).

Visual Evoked Potential (VEP) averages were calculated from the EEG data for each condition. Difference waves of these averages were computed by subtracting No Figure from Figure trials in order to isolate activity related to processing of global organization of the target stimulus and discard processing related to processing of local line elements (Figure 2.2, see also Caputo & Casco, 1999; Lamme, et al., 1992; Scholte, et al., 2006). This was done separately for the Masked and Unmasked condition, as to subtract out direct contributions of the mask to the VEP’s and enable comparison of masked and unmasked trials. All VEP difference waves were converted to spline Laplacian’s (see method section) to be able to
better isolate local generators of any observed effects and filter out deep sources (Nunez & Srinivasan, 2006).

A split half procedure of odd and even trials was used to evaluate the data (Figure 2.3). *Even* trials were used for a visualization of instructive time segments using spline Laplacian maps. These were used to select relevant electrodes for pooling. *Odd* trials were used to evaluate significant deflections of the average current source density at these electrodes over time. Using *odd* trials for the spatial map and *even* trials for the time course values ensures complete independence of data used for inspection and data used for evaluation of significance. This gives an enormous boost to the reliability of the effects that were found, while precluding coincidental significance of sources that were identified visually. It is important to note that even though for brevity not all time points in the 0-305 ms range have been visualized using scalp distribution maps in Figure 2.3, electrode pooling was done on the basis of all visually identifiable sources in this domain, while none were opportunistically left out. Pooling the data to a limited number of electrodes in this way circumvents the multiple comparisons problem of having to evaluate large numbers of electrodes. Pooling relevant electrodes furthermore increases the signal to noise ratio.

Figure 2.3a shows the course of neural processing in the case of unmasked trials. Three stages can be distinguished:

1. A pre-110 ms stage with bilateral anterior occipitotemporal neural generators consistent with feedforward processing towards extrastriate visual areas and beyond (Foxe & Simpson, 2002).
2. A post-110 ms stage with a clear posterior occipital generator consistent with recurrent processing towards early visual areas, possibly V1.
3. A strong recurring bilateral generator in the 200-300 ms range, plausibly extrastriate and beyond.

All generators show significant deflections as ascertained by paired t-tests between Figure and No Figure trials. These were performed on each time point in the time segment, subsequently corrected for multiple comparisons by fixing the False Discovery Rate (FDR) at .05 (see method section).
Figure 2.3 Results. Spline Laplacian maps over three time segments: (1) 78-109 ms, (2) 109-141 ms and (3) 180-305 ms and event related averages of relevant electrodes (pooled occipitally and occipito-temporally). Even trials were used for data inspection (the maps) to determine sensible electrode poolings. Odd trials were used for the averages and evaluation of statistical significance. (a) Unmasked condition (b) Masked condition. Both masked and unmasked show strong significant temporal bilateral activation reflecting feedforward activation in stage 1. More posterior occipital activation reflecting recurrent processing can only be seen in the unmasked condition in stage 2, but is abolished by masking.
Figure 2.3b shows the same three stages for the masked trials. Stage (1) again shows a significant bilateral extrastriate neural generator. Stage (2) and (3) are largely absent and show no significant deflections. A direct comparison of the masked and unmasked difference waves was also done (see Figure 2.4). In accordance with the other results, these showed significant differences in stage (2) and stage (3), but not in stage (1). Taken together, these results are consistent with an interpretation where masking results in an intact initial feedforward activation of extrastriate areas, followed by a disruption of recurrent processing, possibly affecting both striate and extrastriate visual cortex.

Masked vs. Unmasked

![Masked vs. Unmasked](image_url)

Figure 2.4 A difference wave of the difference waves outlined in Figure 2.3 was calculated so as to directly test at which intervals the masked and the unmasked figures differ from each other. This figure clearly shows that masked and unmasked trials do not significantly differ from each other in the early 78-109 ms occipitotemporal stage related to feedforward processing (1), but do differ significantly from each other in the 109-141 ms occipital stage (2) which is plausibly related to recurrent processing as well as the later 180-305 ms occipitotemporal stage (3).

Because the resolution of ERP’s is rather limited, some may object to the idea that recurrent processing can be pinpointed using ERP’s. Yet in our view, only a relatively simple claim needs to be substantiated: (1) there is early activation found anterior on the scalp (2) there is distinct later activation posterior on the scalp. In combination with current source densities, and provided that the generators are clearly distinct, this simple anterior/posterior claim seems sufficient to claim recurrent processing,
Discussion

The most salient aspect of these results is that even when subjects are unable to perceive anything by objective measures, widespread bilateral occipitotemporal activation still takes place, showing a clear ability of the brain to pick up differences that the subject cannot express. This early activation is almost indistinguishable from early activity in the visible situation and is insufficient to generate visual awareness. Some additional process appears necessary in order to achieve conscious perception and detection. The earliest difference between the visible and non-visible condition arises in the >110 ms timeframe, and occurs more posterior than the early bilateral activation. This is most consistent with an interpretation where masking disrupts recurrent processing between high and low visual areas, while leaving feedforward activation intact.

It is unclear how masking accounts resting solely on lateral inhibition could explain these data. The strongest evidence for the influence of lateral inhibition in masking comes from studies employing first order metacontrast stimuli (e.g. Macknik & Livingstone, 1998). First order (luminance defined) stimuli are obvious candidates for spatiotemporal lateral inhibitory influences, since one can see how a metacontrasting stimulus laterally inhibits a first order contrast stimulus of opposite polarity. However, the present study makes use of second order target stimuli and pattern masking. As target and mask do not bear a metacontrasting relation to each other, low level lateral inhibitory influences are less likely. Moreover, any remains of low level inhibitory influences that may still exist are subtracted out using the Figure minus No Figure subtraction.

However, one could postulate that high level lateral inhibitory mechanisms operate between complex representations – as suggested by Macknik and Martinez-Conde (2004). If this were the case, one would predict the pre-110 ms activation to be disrupted in the masked scenario as a result of these inhibitory influences. But the locus of interruption in this experiment is later in time and more posterior. This is not to say that lateral inhibitory influences do not play a role in many types of masking, only that the view of ubiquitous lateral inhibition as an explanation for our results is not tenable.
Interchannel inhibition accounts of masking presume the existence of two channels in visual processing, a fast and a slow one. Masking is presumably caused by the mask in a fast acting channel inhibited by the processing of the target in a slow acting channel. Such accounts cannot be ruled out entirely by these data, as such theories do not specify at which level inhibition takes place. It could be argued that the feedforward activity found in this study is caused by target activity in a fast acting channel, and that the posterior >110 ms activity which we call recurrent activity is caused by a slow acting channel. In the masked scenario this activity would be inhibited by the mask through subsequent activation of the fast acting channel (which would have to be projecting downwards to inhibit the slow acting channel). However, aside from the fact that the posterior activity we observed is not very “sustained” as such models propose, there is little neurophysiological evidence that interchannel inhibition actually exists (also see Enns & Di Lollo, 2000). This makes an explanation built on proven notions of recurrent processing much more parsimonious in this context.

Aside from the present study, a number of other studies have inferred the influence of masking on recurrent processing in human subjects, most notably studies employing object substitution masking (Di Lollo, et al., 2000; Weidner, Shah, & Fink, 2006; Woodman & Luck, 2003). In object substitution masking (also referred to as four-dot masking), a four dot mask is presented surrounding a non-attended target. The subsequent discrimination of this target is unimpaired when target and mask co-terminate, but strongly impaired when the dots remain visible after target offset. This is explained by postulating that the four dots (after offset of the target) create a mismatch between the cortical representation of the four dots in early visual areas and the cortical representation of the target in higher areas. Although we believe the results of these studies have significant implications for the influence of recurrent processing on perception, they differ from our results in important ways.

Four dot masking depends on attention being directed elsewhere and increases with increasing set size (Di Lollo, et al., 2000; Enns, 2004). In our pattern masking study, attention is fully directed on the target, and set size is just one. Moreover, uninterrupted processing of masked targets using object substitution continues well...
Masking disrupts recurrent processing into the 200-300 ms domain (Woodman & Luck, 2003), whereas our results show that processing is disrupted at 110 ms. Indeed, a recent fMRI study comparing pattern and object substitution masking shows that differences between the two can be found in the left middle frontal gyrus, in the precentral gyrus of the right hemisphere and in the medial superior frontal gyrus (Weidner, et al., 2006), while our study shows strictly occipital disruption due to pattern masking.

What could account for these differences? What happens during object substitution masking seems to be more akin to what happens during other attentional manipulations, such as the attentional blink. Many attentional blink studies have shown that processing of unseen target stimuli goes almost completely uninterrupted up to 300 ms and that differences between reported and unreported targets are to be found in frontal and parietal cortices (Kranzioch, Debener, Schwarzbach, Goebel, & Engel, 2005; Luck, Vogel, & Shapiro, 1996; Sergent, Baillet, & Dehaene, 2005). This is reminiscent of the response profile found in object substitution masking. As other authors have noted, the failure of subjects to report stimuli in such instances may reflect a loss of information at a post-perceptual stage, where information is replaced before it can be transferred to working memory and accessible awareness (e.g. Luck, et al., 1996; Woodman & Luck, 2003).

It may still be recurrent processing that is interrupted, but at a later stage and between visual and higher brain areas (possibly frontal and parietal), while early recurrent processing within visual areas goes uninterrupted. Recent models of consciousness proposed by Dehaene, Changeux, Naccache, Sackur and Sergent (2006) and Lamme (2003, 2006) outline similar ideas in more detail. The notion of several reentrant or recurrent loops at multiple levels is highly likely given the temporal dynamics of cortical processing (e.g. Foxe & Simpson, 2002). Because of the +110 ms timing and location observed in this experiment, we conclude that recurrent processing was disrupted at perceptual levels early on, comparable to what has been shown in other studies using different stimuli (Breitmeyer, et al., 2004; Haynes, Driver, & Rees, 2005; Ro, et al., 2003).

The function of recurrent processing within occipital cortex is a somewhat open question, though strong evidence exists that it is of importance for figure-ground segmentation (e.g. Hupe, et al., 1998;
Lamme, 1995; Lamme, Rodriguez-Rodriguez, & Spekreijse, 1999). In such studies it is typically shown that a neuron in V1 starts responding differentially after about 100 ms when its classical receptive field (CRF) is located on the center of a figure compared to when it is on a background, despite local stimulation being exactly equal (cf. Figure 2.2). These extra-classical receptive field effects are usually referred to as contextual modulation, as they are modulated by context information outside their CRF. The presence and timing (>100 ms) of contextual modulation strongly suggests that recurrent neural processing takes place going back from higher areas containing larger receptive fields to lower areas such as V1. Also, when extrastriate regions are lesioned, contextual modulation disappears, while leaving receptive field tuning properties intact, conclusively showing the role of recurrent processing in contextual modulation (Lamme, Super, et al., 1998).

A study in which the effect of target-mask SOA on contextual modulation in macaque V1 was investigated showed a strong relation between SOA, contextual modulation, and the monkey’s ability to detect a texture defined figure (Lamme, et al., 2002). Moreover, any influence of SOA on the CRF tuning properties of these neurons was almost completely absent, showing feedforward activation was largely preserved. However, since measurements were done only in V1, it was unclear to what extent feedforward activation would continue to higher areas and to what extent the same would hold for humans.

The present study clearly demonstrates that in humans, undetected and strongly masked texture stimuli are processed far beyond striate cortex (see Figure 2.3), showing that feedforward processing is preserved, while recurrent processing is selectively interrupted. A neural network model of figure-ground segregation by Roelfsema, Lamme, Spekreijse and Bosch (2002) in which texture segregation in macaques is modeled in terms of feedforward and recurrent processing serves extremely well in explaining the spatio-temporal profile of such results.

Related models propose that recurrent processing is more intimately related to visual awareness (e.g. Lamme, 2001; Ro, et al., 2003). These views on recurrent processing are not necessarily contradicting, as visual awareness might simply be the way surface
segregation and related processes express themselves phenomenologically. The view that recurrent activity in visual cortex correlates with visual awareness is now supported by converging evidence from monkey physiology (e.g. Lamme, Super, Landman, Roelfsema, & Spekreijse, 2000), EEG (the present study), TMS (Pascual-Leone & Walsh, 2001) and fMRI (e.g. Haynes, et al., 2005).

**Conclusion**

We argue and found evidence for at least three stages in early visual processing: (1) a completely unconscious pre-110 ms feedforward stage which is probably involved in boundary and object detection but goes uninterrupted by masking (2) a second stage, which operates in the 110-140 ms range reflecting recurrent processing in visual cortex, plausibly encapsulating processes such as figure-ground segmentation and phenomenal awareness and (3) a stage in the 200-300 ms range which is contingent upon earlier stages. Specifically the second stage was interrupted by backward masking, confirming that pre-110 ms activation proceeds outside awareness, whereas more posterior >110 ms activity is (at least) a necessary condition for awareness to arise. Finally, we reject explanations of backward masking as a unitary phenomenon, in agreement with other authors such as Bachmann et al. (2005). Any explanation of the masking phenomenon should carefully take into account physical stimulus characteristics and how these are processed by the various components of the visual system.